

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number

TO: Ben Sackey

Location: 5b31 / 5c18

Art Unit: 1626

Friday, January 06, 2006

Case Serial Number: 10 / 705659

From: Noble Jarrell

Location: Biotech-Chem Library

Rem 1B71

Phone: 272-2556

Noble.jarrell@uspto.gov

Search Notes	
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175358

Nable

Scientific and Technical Information Center

SEARCH REQUEST FORM

	SEARCH REQU	EST FORM	
Requester's Full Name: BE Art Unit: 1636 Phot Location (Bldy/Room#): BE 55	ne Number: 2- 0704 83/Mailbox #): Res	Serial Number: 10/2	os, 659 PAPER DISK
To ensure an efficient and quality scare			
Title of Invention: Campo	mals and Synth	esis process.	
Title of Invention: <u>Compo</u> Inventors (please provide full names	s): milliam J.	Segrey	
Earliest Priority Date: 11	03		
Search Topic: Please provide a detailed statement of the elected species or structures, keywords, sy Define any terms that may have a special	nonyms, acronyms, and registry num	bers, and combine with the concept or i	searched. Include the willing of the invention.
For Sequence Searches Only Fleave in appropriate serial number.	clude all pertinent information (pare	nt, child, divisional, or issued patent nu	mbers) along with the
A process by the not compris	~ preparing	6-chloro - 2,5	-dica bonamida
Thenal campris	ing chlorinal	- 2- alkyl-6	then 0 : 1
2- alkyl	-6-anino-7-c	May 0 star 10 yr 30	,
		(STIC)	7,000 A
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STAFF USE ONLY	Type of Search	Vendors and cost where app	licable
iearcher: Able	NA Sequence (#)	STN STN	_Dialog
Searcher Phona #:	AA Sequence (#)	Questel/Orbit	Lexis/Nexis
earcher Location:	5 Structure (#)	Westlaw	
Date Searcher Picked Up: 11606	Bibliographic	In-house sequence system	
Date Completed: 1666	Litigation	CommercialOligomer	Score/Length
earcher Prep & Review Time: 10	Fulltext	Interference SPDI Other (specify)	Encode/Transl

Other

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STRUCTURE FILE UPDATES: 4 JAN 2006 HIGHEST RN 871209-00-6
DICTIONARY FILE UPDATES: 4 JAN 2006 HIGHEST RN 871209-00-6

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

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Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

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=> d que sta 110 L5 STR 11 N 2 7 Ak 10 1 C 3 O 8 6 C C 8

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 11

STEREO ATTRIBUTES: NONE L10 632 SEA FILE=REGISTRY SSS FUL L5

100.0% PROCESSED 4059 ITERATIONS SEARCH TIME: 00.00.01

632 ANSWERS

=> d que sta l13

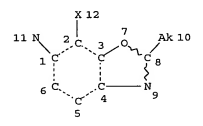
NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 11

STEREO ATTRIBUTES: NONE

L10 632 SEA FILE=REGISTRY SSS FUL L5

L11 STR



NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 12

STEREO ATTRIBUTES: NONE

L13 7 SEA FILE=REGISTRY SUB=L10 SSS FUL L11

100.0% PROCESSED 126 ITERATIONS 7 ANSWERS

SEARCH TIME: 00.00.01

=> d que sta 116

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 14

STEREO ATTRIBUTES: NONE

68 SEA FILE=REGISTRY SSS FUL L14

100.0% PROCESSED 3134 ITERATIONS 68 ANSWERS

SEARCH TIME: 00.00.01

=> b hcap FILE 'HCAPLUS' ENTERED AT 08:52:55 ON 06 JAN 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 6 Jan 2006 VOL 144 ISS 2 FILE LAST UPDATED: 4 Jan 2006 (20060104/ED) FILE LAST UPDATED: 4 Jan 2006

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d all hitstr 124 tot

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L24 ANSWER 1 OF 3 HCAPLUS COPYRIGHT 2006 ACS on STN
    2005:411078 HCAPLUS
AN
DN
     142:463458
     Entered STN: 13 May 2005
ED
     Process for preparing 6-chloro-2,5-dicarbonamidophenol compounds
TT
    Begley, William J.
IN
PA
     Eastman Kodak Company, USA
```

SO U.S. Pat. Appl. Publ., 9 pp.

CODEN: USXXCO

DТ Patent LΑ

English

ICM C07D-0263/52 ICS C07C-0231/10

INCL 548217000; 564155000

25-19 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)

PAN	.CNI I																
	PATENT	NO.			KIN)	DATE		7	APPL:	ICAT:	I NOI	NO.		D2	ATE	
												- 					
ΡI	US2005101784			A1 20050512			2003US-0705659						20031110				
	WO2005047271				A1 20050526			2004WO-US36261						20041029			
	W:	AE,	AG,	AL,	AM,	AT,	ΑU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,
							DE,										
							ID,										
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NΑ,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
							TZ,										
	RW:						MW,										
							RU,										

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EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
             SN, TD, TG
PRAI 2003US-0705659
                                 20031110
CLASS
                 CLASS PATENT FAMILY CLASSIFICATION CODES
PATENT NO.
                        C07D-0263/52
US 2005101784
                 ICM
                        C07C-0231/10
                 ICS
                 INCL
                        548217000; 564155000
                 IPCI
                        C07D0263-52 [ICM, 7]; C07C0231-10 [ICS, 7]
                 NCL
                        548/217.000
                 ECLA
                        C07C315/04; C07D263/56B
                        C07D0263-56 [ICM, 7]; C07C0317-22 [ICS, 7]
                 IPCI
WO2005047271
                        C07C315/04; C07D263/56B
                 ECLA
     Disclosed is a process for preparing a 6-chloro-2,5-dicarbonamidophenol
     compds. comprising a step employing a 2-alkyl-6-aminobenzoxazole to form a
     2-alkyl-6-amino-7-chlorobenzoxazole in which the 2-alkyl group is
     unbranched at the \alpha-carbon. It also provides intermediate compds.
     useful in the process. The process provides a simple and safe way to
     prepare 6-chloro-2,5-dicarbonamidophenol compds. in good yield. Thus,
     nitration of 5-chloro-2-methylbenzoxazole by HNO3/H2SO4 at 20° gave
     5-chloro-2-methyl-6-nitrobenzoxazole which was reduced over Raney nickel
     in THF at room temperature under H pressure of 50 psi to give
     6-Amino-5-chloro-2-methylbenzoxazole (I). Chlorination of I by sulfuryl
     chloride in EtOAc for 1 h gave 6-amino-5,7-dichloro-2-methylbenzoxazole
     which was acylated by 2-[(4-dodecyloxyphenyl)sulfonyl]butanoyl chloride in
     pyridine/EtOAc/THF at 15° for 30 min to give 6-[[2-[(4-
     dodecyloxyphenyl) sulfonyl] butanoyl] amino] -5,7-dichloro-2-methylbenzoxazole
     (II). Hydrolysis of II in a mixture of concentrated HCl and THF at 65° for
     .apprx.3 h gave 6-amino-2,4-dichloro-3-[[2-[(4-
     dodecyloxyphenyl)sulfonyl]butanoyl]amino]phenol which was acylated by
     3,4-dichlorobenzoyl chloride in pyridine/THF at room temperature for 30 min to
     give 2,4-dichloro-3-[[2-[(4-dodecyloxyphenyl)sulfonyl]butanoyl]amino]-6-
     (3,4-benzoylamino)phenol.
st
     alkylaminochlorobenzoxazole prepn intermediate chlorodicarbonamidophenol;
     chlorodicarbonamidophenol prepn
TT
     701-16-6, 5-Fluoro-2-methylbenzoxazole
                                               3024-72-4, 3,4-Dichlorobenzoyl
               3282-30-2, Pivaloyl chloride 19219-99-9, 5-Chloro-2-
     methylbenzoxazole
                        851486-98-1, 2-[(4-Dodecyloxyphenyl)sulfonyl]butanoic
     acid
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (process for preparing 6-chloro-2,5-dicarbonamidophenol compds. by
        N-acylation of alkyl(amino)benzoxazole derivs. followed by hydrolysis
        and N-acylation)
IT
     13452-16-9P, 5-Chloro-2-methyl-6-nitrobenzoxazole
     40703-40-0P, 5-Fluoro-2-methyl-6-nitrobenzoxazole
                                                          98121-18-7P,
     2-[(4-Dodecyloxyphenyl)sulfonyl]butanoyl chloride 323579-00-6P,
     6-Amino-5-chloro-2-methylbenzoxazole 851486-89-0P,
     6-Amino-5,7-dichloro-2-methylbenzoxazole 851486-90-3P,
     6-[[2-[(4-Dodecyloxyphenyl)sulfonyl]butanoyl]amino]-5,7-dichloro-2-
     methylbenzoxazole
                        851486-91-4P, 2,4-Dichloro-3-[[2-[(4-
     dodecyloxyphenyl) sulfonyl] butanoyl] amino] -6-aminophenol
     851486-92-5P, 2,4-Dichloro-3-[[2-[(4-dodecyloxyphenyl)sulfonyl]but
     anoyl]amino]-6-(3,4-dichlorobenzoylamino)phenol 851486-93-6P,
     6-Amino-5-fluoro-2-methylbenzoxazole 851486-94-7P,
     6-Amino-7-chloro-5-fluoro-2-methylbenzoxazole 851486-95-8P,
     6-[[2-[(4-Dodecyloxyphenyl)sulfonyl]butanoyl]amino]-7-chloro-5-fluoro-2-
                        851486-96-9P, 6-Amino-2-chloro-3-[[2-[(4-
     methylbenzoxazole
     dodecyloxyphenyl) sulfonyl] butanoyl] amino] -4-fluorophenol
     851486-97-0P, 2-Chloro-3-[[2-[(4-dodecyloxyphenyl)sulfonyl]butanoy
     1] amino] -4-fluoro-6-(pivaloylamino) phenol
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (process for preparing 6-chloro-2,5-dicarbonamidophenol compds. by
        N-acylation of alkyl(amino)benzoxazole derivs. followed by hydrolysis
```

and N-acylation) IT 13452-16-9P, 5-Chloro-2-methyl-6-nitrobenzoxazole 40703-40-0P, 5-Fluoro-2-methyl-6-nitrobenzoxazole 323579-00-6P, 6-Amino-5-chloro-2-methylbenzoxazole 851486-89-0P, 6-Amino-5,7-dichloro-2-methylbenzoxazole 851486-90-3P, 6-[[2-[(4-Dodecyloxyphenyl)sulfonyl]butanoyl]amino]-5,7-dichloro-2-methylbenzoxazole 851486-92-5P, 2,4-Dichloro-3-[[2-[(4-dodecyloxyphenyl)sulfonyl]butanoyl]amino]-6-(3,4dichlorobenzoylamino) phenol 851486-93-6P, 6-Amino-5-fluoro-2methylbenzoxazole 851486-94-7P, 6-Amino-7-chloro-5-fluoro-2methylbenzoxazole 851486-95-8P, 6-[[2-[(4-Dodecyloxyphenyl) sulfonyl] butanoyl] amino] -7-chloro-5-fluoro-2methylbenzoxazole 851486-97-0P, 2-Chloro-3-[[2-[(4dodecyloxyphenyl)sulfonyl]butanoyl]amino]-4-fluoro-6-(pivaloylamino)phenol RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (process for preparing 6-chloro-2,5-dicarbonamidophenol compds. by N-acylation of alkyl(amino)benzoxazole derivs. followed by hydrolysis and N-acylation) RN 13452-16-9 HCAPLUS CN Benzoxazole, 5-chloro-2-methyl-6-nitro- (8CI, 9CI) (CA INDEX NAME)

RN 40703-40-0 HCAPLUS
CN Benzoxazole, 5-fluoro-2-methyl-6-nitro- (9CI) (CA INDEX NAME)

$$\bigcap_{O_2N} \bigcap_{O_2N} \bigcap$$

RN 323579-00-6 HCAPLUS
CN 6-Benzoxazolamine, 5-chloro-2-methyl- (9CI) (CA INDEX NAME)

RN 851486-89-0 HCAPLUS CN 6-Benzoxazolamine, 5,7-dichloro-2-methyl- (9CI) (CA INDEX NAME)

$$H_2N$$
 O
 Me

RN 851486-90-3 HCAPLUS
CN Butanamide, N-(5,7-dichloro-2-methyl-6-benzoxazolyl)-2-[[4-

(dodecyloxy) phenyl] sulfonyl] - (9CI) (CA INDEX NAME)

RN 851486-92-5 HCAPLUS

CN Benzamide, 3,4-dichloro-N-[3,5-dichloro-4-[[2-[[4-(dodecyloxy)phenyl]sulfonyl]-1-oxobutyl]amino]-2-hydroxyphenyl]- (9CI) (CA INDEX NAME)

RN 851486-93-6 HCAPLUS

CN 6-Benzoxazolamine, 5-fluoro-2-methyl- (9CI) (CA INDEX NAME)

$$H_2N$$
 N Me

RN 851486-94-7 HCAPLUS

CN 6-Benzoxazolamine, 7-chloro-5-fluoro-2-methyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{C1} & \text{Me} \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ \end{array}$$

RN 851486-95-8 HCAPLUS

CN Butanamide, N-(7-chloro-5-fluoro-2-methyl-6-benzoxazolyl)-2-[[4-(dodecyloxy)phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

RN 851486-97-0 HCAPLUS

CN Butanamide, N-[2-chloro-4-[(2,2-dimethyl-1-oxopropyl)amino]-6-fluoro-3-hydroxyphenyl]-2-[[4-(dodecyloxy)phenyl]sulfonyl]- (9CI) (CA INDEX NAME)

```
L24
    ANSWER 2 OF 3 HCAPLUS COPYRIGHT 2006 ACS on STN
AN
     1967:2504 HCAPLUS
DN
     66:2504
     Entered STN: 12 May 1984
ED
     Syntheses of heterocyclic compounds. XIV. Oxazoles from the pyrolysis of
ΤI
     aryl azides in a mixture of a carboxylic and polyphosphoric acid
     Garner, Robert; Mullock, E. B.; Suschitzky, Hans
ΑU
CS
     Roy. Coll. Advan. Technol., Salford, UK
     Journal of the Chemical Society [Section] C: Organic (1966), (21), 1980-3
SO
     CODEN: JSOOAX; ISSN: 0022-4952
DT
     Journal
     English
LΑ
CC
     28 (Heterocyclic Compounds (More Than One Hetero Atom))
     For diagram(s), see printed CA Issue.
GΙ
     cf. CA 65, 15366b. Aromatic azides (I) with a para-substituent decompose
AB
     thermally in a mixture of polyphosphoric and a carboxylic acid to give
     oxazoles (II), or in some cases N,O-diacyl o-aminophenols, in good yield.
     Various aspects of this nitrene mechanism are discussed. 18 references.
ST
     OXAZOLES BENZO; AZIDES; BENZOXAZOLES
IT
     Aryl azides
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (pyrolysis of)
IT
     288-42-6D, Oxazole, derivs.
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (from azide pyrolysis)
     833-62-5P 5683-43-2P 13243-31-7P
                                          13243-32-8P
                                                        13243-36-2P
IT
                                          13243-40-8P
     13243-37-3P 13243-38-4P 13243-39-5P
                                 13452-14-7P
                                               13452-15-8P 13452-16-9P
     13438-55-6P
                   13452-13-6P
                                 14724-89-1P
     13452-17-0P
                   13473-67-1P
                                               15260-89-6P
     34594-87-1P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
     5683-43-2P 13243-38-4P 13243-39-5P
     13452-16-9P 13452-17-0P
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (preparation of)
RN
     5683-43-2 HCAPLUS
     Benzoxazole, 2-methyl-6-nitro- (6CI, 7CI, 8CI, 9CI) (CA INDEX NAME)
CN
```

RN 13243-38-4 HCAPLUS CN Benzoxazole, 7-chloro-2-methyl-6-nitro- (8CI) (CA INDEX NAME)

RN 13243-39-5 HCAPLUS

CN Benzoxazole, 2-ethyl-6-nitro- (8CI) (CA INDEX NAME)

13452-16-9 HCAPLUS RN

Benzoxazole, 5-chloro-2-methyl-6-nitro- (8CI, 9CI) (CA INDEX NAME) CN

13452-17-0 HCAPLUS RN

CN Benzoxazole, 2,2'-octamethylenebis[6-nitro- (8CI) (CA INDEX NAME)

$$N$$
 $O2N$ $O2N$

L24 ANSWER 3 OF 3 HCAPLUS COPYRIGHT 2006 ACS on STN

AN 1953:41191 HCAPLUS

DN 47:41191

OREF 47:6894f-i,6895a-e

ED Entered STN: 22 Apr 2001

Quinone imides. XVIII. p-Quinonedipivalimides and their reactions ΤI

ΑU Adams, Roger; Stewart, John M.

CS Univ. of Illinois, Urbana

Journal of the American Chemical Society (1952), 74, 3660-4 SO

CODEN: JACSAT; ISSN: 0002-7863

DTJournal

LΑ Unavailable

CC 10 (Organic Chemistry)

AΒ To 10.8 g. p-C6H4(NH2)2 (I) in 125 cc. pyridine was added slowly with stirring 25.3 g. Me3CCOCl (II) (obtained nearly quantitatively by refluxing the acid 1 hr. with excess SOCl2), and the mixture poured after 6 hrs. into excess HCl and ice to give 25.5 g. (93%) p-C6H4(NHCOCMe3)2, m. 283° (from dioxane). Similarly was prepared from 4.6 g. Ph3CCOCl (obtained by carbonation of Ph3CMgCl and treatment of the acid with SOCl2) in 20 cc. pyridine and 0.81 g. I in 10 cc. pyridine 4.5 g. (92.5%) p-C6H4(NHCOCPh3)2, m. 324-6° (from HCONMe2). I (5.0 g.) and 20 cc. CF3CO2H refluxed 7 hrs., and the mixture poured into 400 cc. dilute HCl

yielded 5.9 q. (38%) p-C6H4(NHCOCF3)2, m. 274° (from dioxane), which, refluxed with Pb(OAc)4 in CHCl3, tarry, amorphous products. A similar oxidation of p-C6H4(NH-COCCl3)2 gave a small yield of an unstable product. Equimolar amts. of N,N'-(p-phenylene)dipivalamides (III) and Pb(OAc)4 refluxed 2 hrs. in dry CCl4 (25 cc./g. III) yielded the corresponding substituted N, N'-dipivalyl-p-quinone diimines [substituent, m.p., % yield given]: H (V), 164.5°, 84; 2-Cl (VI), 81-8.5°, 65; 2,6-Cl2 (VII), 104.5-6.5°, 85; 2,5-Cl2 (VIII), 159.5-60.5°, 80; 2,3-Cl2 (IX), 138.5-9.5°, 75; 2,3,5-Cl3 (X), 115.5°, 78 (all recrystd. from petr. ether), p-C6H4(NHCOCMe3)2 (0.83 g.) and an equivalent amount of Pb(OAc)4 in Ac2O stirred 24 hrs. at 70° and the mix decomposed with 500 cc. H2O gave 31% 2,3,5,6-tetrachloro-IV, m. 200.5-1°. HCl passed into a petr. ether solution of the IV precipitated Cl-substituted III (substituent, m.p., and yield given): 2-Cl, 215°, 90; 2,6-Cl2 (XI), 257°, 95, from VI; 2,3,5-Cl3 (XII), m. 205-6° with rapid shrinking at 196°, 96% from VIII, 97% from IX, and 24% from VII (all from CHCl3-petr. ether); and 2,3,5,6-Cl4 (XIII), m. 335-5.5° (from CHCl3), 44 yield together with 45% 2-tert-butyl-4,5,7-trichloro-6-(pivalylamino)benzoxazole, m. 225° (from aqueous EtOH), from X. V (0.69 g.) in 10 cc. glacial AcOH let stand 1 day at room temperature and poured into H2O yielded 0.3 g. (36%) 2,1,4-AcOC6H3(NHCOCMe3)2 (XIV), needles, m. 156.5-7° (from CHCl3). To 9.0 g. V added slowly with stirring and cooling 9 cc. 98% HCO2H, and the mixture diluted with cold Et2O yielded 8.5 g. (81%) 2,1,4-HCOC6H3(NH-COCMe3)2 (XV), platelets, m. 200-1° (from CHCl3). XIV refluxed 30 min. with 10% aqueous NaOH gave 2,5-(Me3CCONH)2C6H3OH (XVI), needles, m. 248° (from petr. ether), also obtained in 90% yield by boiling 4.0 g. XV in 100 cc. (CH2OH)2 5 min., or in 77% yield by alkaline hydrolysis of XV. XVI (0.30 g.) heated 15 min. at 250° and 100 mm. pressure and the cooled melt triturated with 15 cc. Et2O yielded 0.18 g. (84%) 2-tert-butyl-6-(pivaloylamino)benzoxazole, platelets or needles, m. 164-5° (from Et2O-petr. ether), hydrolyzed to XVI by refluxing 3 hrs. with 10% aqueous NaOH. The following Cl-substituted III (substituent given) were prepared from the corresponding Cl-substituted 1.2HCl salts and II in pyridine: 2,5-Cl2, needles, m. 239-40° (from CHCl3-petr. ether); 2,3-Cl2, needles, 61%, m. 200-1° (from MeOH) [the free diamine, needles, m. 120.5-1° (from H2O)]; XI, m. 256-7°, in poor yield. HCl passed 10 min. into 7.0 g. VII in 250 cc. petr. ether and the product chromatographed from 8 l. 3:1 petr. ether-Et20 mixture on activated Al2O3 gave 24% XII and 33% 2-tert-butyl-5,7-dichloro-6-(pivalylamino)benzoxazole (XVII), needles, m. 165.5-7.5° (from petr. ether). Aqueous alkaline hydrolysis of XVII yielded 88% 2,4,3,6-Cl2(Me3CCONH)2C6HOH (XVIII), needles, m. 226° (from Et20-petr. ether). XVIII in (CH2OH)2 boiled 10 min. gave XVII. Cl passed into 2.0 g. XVI in 100 cc. glacial AcOH at 20° to a weight increase of 0.95 g., and the mixture E poured into 600 cc. cold H2O gave XVIII. To 0.12 g. XVIII in 20 cc. H2O and 0.5 cc. 5% aqueous NaOH was added with stirring 0.04 cc. Ac2O to give 0.10 g. (83%) acetate of XVIII, m. 267-8° (from CHCl3-petr. ether). IX (0.3 g.) in 5 cc. 98% HCO2H let stand 0.5 hr. at room temperature, the red solution diluted with 75 cc. Et20, extracted with 100 cc. 5% aqueous NaOH, and the extract acidified with HCl yielded 0.2 g. (63%) 3,4,2,5-Cl2(Me3CCONH)2C6HOH, m. 190.5-1° (from CHCl3-petr. ether). Similarly was prepared 3,6,2,5-Cl2(Me3CCONH)2C6HOH, 44%, m. 199.5-200.5° (from CHCl3-petr. ether), from VIII. p-C6Cl4(NH2)2 (2.5 g.) and 2.5 g. II in 25 cc. pyridine refluxed 4.5 hrs. gave 4.0 g. (95%) XIII. Quinone imines Oxidation (of N, N'-p-phenylenebis amides) Propionamide, N,N'-[2,3-dichloro-3-5-hydroxy-p-phenylene)bis[2,2-dimethyl-Propionamide, N,N'-[2,3-dichloro-5-5-hydroxy-p-phenylene)bis[2,2-dimethyl-Propionamide, N, N'-[2,5-dichloro-3-5-hydroxy-p-phenylene)bis[2,2-dimethyl-Propionamide, N,N'-[2,5-dichloro-5-5-hydroxy-p-phenylene)bis[2,2-dimethyl-859057-55-9, Propionamide, N,N'-(hydroxy-p-phenylene)bis[2,2-dimethyl-(and esters)

4257-74-3, Acetamide, N,N'-p-phenylenebis[2,2,2-trichloro-

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(oxidation of) TT 404-28-4, Acetamide, N,N'-p-phenylenebis[2,2,2-trifluoro-6068-70-8, Acetyl chloride, triphenyl- 6937-98-0, Propionamide, N, N'-p-phenylenebis[2,2-dimethyl- 41946-53-6, p-Phenylenediamine, 2,3-dichloro- 313394-34-2, Propionamide, N,N'-(chloro-pphenylene)bis[2,2-dimethyl-854052-01-0, p-Benzoquinone diimine, 2,3,5-trichloro-N,N'-dipivaloyl- 854052-02-1, p-Benzoquinone diimine, 2,3,5,6-tetrachloro-N,N'-dipivaloyl- 854052-25-8, p-Benzoquinone diimine, 2-chloro-N, N'-dipivaloyl- 854053-46-6, p-Benzoquinone diimine, 2,6-dichloro-N,N'-dipivaloyl- 854053-47-7, p-Benzoquinone diimine, 2,5-dichloro-N,N'-dipivaloyl-854053-48-8, p-Benzoquinone diimine, 2,3-dichloro-N,N'-dipivaloyl- 854164-19-5, Benzoxazole, 2-tert-butyl-4,5,7-trichloro-6-pivalamido- 854164-19-5, Propionamide, N-(2-tert-butyl-4,5,7-trichloro-6-benzoxazolyl)-2,2-dimethyl-854164-20-8, Benzoxazole, 2-tert-butyl-6-pivalamido-854164-20-8, Propionamide, N-2-tert-butyl-6-benzoxazolyl-2,2dimethyl- 854164-22-0, Benzoxazole, 2-tert-butyl-5,7-dichloro-6pivalamido- 854164-22-0, Propionamide, N-(2-tert-butyl-5,7dichloro-6-benzoxazolyl)-2,2-dimethyl- 855464-56-1, p-Benzoquinone diimine, N,N'-dipivaloyl- 856985-71-2, Propionamide, N, N'-(2,6-dichloro-p-phenylene) bis [2,2-dimethyl-856985-73-4, Propionamide, N,N'-[2,5-dichloro-p-phenylene]bis[2,2-dimethyl-856985-75-6, Propionamide, N,N'-[2,3-dichloro-p-phenylene]bis[2,2-dimethyl-·856985-79-0, Propionamide, N, N'-(3,5-dichloro-2-hydroxy-pphenylene)bis[2,2-dimethyl-, acetate 857231-91-5, Propionamide, N, N'-(trichloro-p-phenylene)bis[2,2-dimethyl- 857943-06-7, Propionamide, N, N'-(tetrachloro-p-phenylene)bis[2,2-dimethyl-859301-32-9, Propionamide, N,N'-(3,5-dichloro-2-hydroxy-p-phenylene)bis[2,2-dimethyl-861058-90-4, Acetamide, N,N'-p-phenylenebis[2,2,2-triphenyl-(preparation of) TT 854164-19-5, Benzoxazole, 2-tert-butyl-4,5,7-trichloro-6pivalamido- 854164-20-8, Benzoxazole, 2-tert-butyl-6-pivalamido-854164-22-0, Benzoxazole, 2-tert-butyl-5,7-dichloro-6-pivalamido-859301-32-9, Propionamide, N, N'-(3, 5-dichloro-2-hydroxy-pphenylene) bis [2, 2-dimethyl-(preparation of) RN854164-19-5 HCAPLUS CN INDEX NAME NOT YET ASSIGNED

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RN 859301-32-9 HCAPLUS

CN Propionamide, N,N'-(3,5-dichloro-2-hydroxy-p-phenylene)bis[2,2-dimethyl-(5CI) (CA INDEX NAME)

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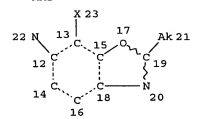
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